

Role of Transcription Factor Kar4 in Regulating Downstream Events in the *Saccharomyces cerevisiae* Pheromone Response Pathway[∇]

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Yeast Kar4 is a putative transcription factor required for karyogamy (the fusion of haploid nuclei during mating) and possibly other functions. Previously known to be required only for the transcriptional induction of *KAR3* and *CIK1*, microarray experiments identified many genes regulated by Kar4 in both mating and mitosis. Several gene clusters are positively or negatively regulated by mating pheromone in a Kar4-dependent manner. Chromatin immunoprecipitation and gel shift assays confirmed that Kar4 binds to regulatory DNA sequences upstream of *KAR3*. Together with one-hybrid experiments, these data support a model in which both Kar4 and Ste12 bind jointly to the *KAR3* promoter. Analysis of the upstream regions of Kar4-induced genes identified a DNA sequence motif that may be a binding site for Kar4. Mutation within the motif upstream of *KAR3* eliminated pheromone induction. Genes regulated by Kar4, on average, are delayed in their temporal expression and exhibit a more stringent dose response to pheromone. Furthermore, the induction of Kar4 by pheromone is necessary for the delayed temporal induction of *KAR3* and *PRM2*, genes required for efficient nuclear fusion during mating. Accordingly, we propose that Kar4 plays a critical role in the choreography of the mating response.

In the budding yeast *Saccharomyces cerevisiae*, mating occurs between two haploid cells of different mating types, *MATa* and *MAT α* (reviewed in references 9 and 31). Mating is initiated by the exchange of specific mating pheromones; *MATa* cells secrete *a*-factor and *MAT α* cells secrete α -factor. Binding of the pheromone to its specific transmembrane receptor elicits a complex signal transduction pathway, resulting in cell cycle arrest and the transcriptional induction of mating-specific genes (reviewed in references 11 and 47). Mating-induced transcription requires activation of the transcription factor Ste12, which binds to pheromone response elements (PREs) in the upstream activating sequences of target genes (12, 16). Ste12 is also involved in the regulation of genes required for a distinct developmental program: filamentous growth in response to starvation (28).

Exposure of cells to mating pheromone leads to the induction of genes that are regulated by Ste12 alone (16), as well as genes that require a second transcription factor, Mcm1 (35, 50). Similarly, during filamentous growth, Ste12-dependent induction requires Tec1, with which it binds cooperatively at filamentation-responsive elements (2, 20, 27, 50). Thus, differential gene expression by Ste12 is achieved through cooperative interactions with other transcription factors such as Tec1 and Mcm1 and possibly others.

Mating consists of the fusion of two haploid cells, followed by the congression and eventual fusion of the two nuclei in an event called karyogamy (reviewed in reference 41). Preceding cell fusion, cytoplasmic microtubules stemming from the spindle pole body move each nucleus to the mating projection of

each haploid cell (26, 34). Upon cell fusion, the cytoplasmic contents of each cell intermingle, facilitating the connection of cytoplasmic microtubules via Kar3, a kinesin-related protein, and Cik1, its associated light chain (33, 38). During nuclear congression, movement and microtubule shortening are produced by Kar3 and Cik1 acting on the cytoplasmic microtubules (25, 26, 33). Finally, the nuclear envelopes fuse at the spindle pole bodies, resulting in the formation of a single diploid nucleus (3).

The mating process must be precisely regulated both spatially and temporally. For example, premature cell wall degradation prior to cell adhesion, at ectopic or supernumerary sites, would be catastrophic. Moreover, many of the genes induced during mating, such as Kar3, have other functions during mitosis, and their premature overexpression may be deleterious during the cell cycle. The current model for the pathway of pheromone-induction suggests that all mating-induced genes are regulated by Ste12, providing no mechanism for the sequential expression of genes during mating.

Kar4 is a good candidate for controlling the temporal regulation of genes that show a delay during the pheromone response. Kar4 was identified previously as a putative transcription factor, which is required, together with Ste12, for the transcriptional induction of *KAR3* and *CIK1* by pheromone (14, 23). Previous work demonstrated that a 30-bp DNA sequence upstream of *KAR3* is necessary for Kar4- and Ste12-dependent induction by pheromone (23). Overexpression of Ste12 is sufficient to induce the expression of *KAR3*, in the absence of both Kar4 and pheromone (23). In contrast, the overexpression of Kar4 does not bypass the requirement for Ste12, although it does partially alleviate the requirement for pheromone (23). These data suggest that Kar4 facilitates Ste12's transcriptional induction of genes such as *KAR3*.

Together with yeast Ime4, Kar4 is a member of the large MT-A70 superfamily of proteins, which includes the *S*-adeno-

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sylmethionine binding subunit of mRNA:m⁶A methyl transferases (MTase) and bacterial DNA:m⁶A MTases (5). Ime4 is required for the appearance of m⁶A in mRNA during yeast sporulation (7). In contrast, Kar4 is the prototypical member of highly divergent MT-A70 subfamily, conserved within fungi, in which the critical residues in the S-adenosylmethionine binding site have been altered (5). Kar4 and other members of this subfamily are unlikely to be MTases (5).

The regulation of Kar4 expression is highly complex, occurring at the level of transcription, translation, and protein turnover. There are two forms of Kar4: a constitutive, low-abundance 38.5-kDa protein (Kar4-long), which predominates during vegetative growth, and an abundant 35.5-kDa protein, which is induced by mating pheromone (Kar4-short) (14, 23). The two forms differ at their amino termini; Kar4-short is translated from a shorter mRNA and initiated from an AUG codon that is internally encoded in the Kar4-long message. In addition to differential transcriptional regulation, the two protein forms have different rates of protein synthesis and turnover (14).

The induction of *KAR4* transcription by Ste12 after exposure to pheromone potentially provides an additional level of regulation for a subset of genes expressed during mating. In particular, the requirement that Kar4 act along with Ste12 for the induction of *KAR3* and *CIK1* might provide a mechanism for temporal regulation (23). Formally, Kar4 and Ste12 comprise a “feed-forward loop” regulatory circuit (29, 30). In this model, Kar4 and Ste12 jointly regulate the transcription of their target genes, providing a delayed response to the inducing signal.

The complexity of its regulation underscores the likelihood that Kar4 performs other regulatory functions beyond the mating-dependent regulation of *KAR3* and *CIK1*. In support of this hypothesis, Kar4 is required for meiosis, its overexpression is inhibitory to vegetative growth, and *kar4* deletion causes a cell cycle delay in G₁ (23). Consistent with multiple functions for Kar4, we show here that Kar4 regulates a large array of genes in both mating and mitosis. Remarkably, the set of genes affected by Kar4 includes both positively and negatively regulated genes.

MATERIALS AND METHODS

Strains and microbial techniques. The yeast strains and plasmids used in the present study are listed in Table 1. Standard yeast media and genetic techniques were as described previously (42). Synthetic alpha factor (Princeton Syn/Seq facility) was used at 3 to 10 μM. Microscopic analysis of karyogamy in zygotes by DAPI (4',6'-diamidino-2-phenylindole) staining was performed as described previously (22). To analyze the defect of *kar4*^{C214G/C217G}, matings of host *kar4Δ* strains MS3216 and MS3212 were performed. Each strain harbored either the wild-type *KAR4* plasmid pMR2654, the *kar4*^{C214G/C217G} plasmid pMR5511, or the vector pRS416 (45). Analysis of the *prm2* mating defect was performed by crossing *MATa* (Y01430) and *MATα* (Y11430) deletion strains; wild-type control strains were MS3265 and MS3213 and the *kar4Δ* strains were MS3216 and MS3212. Matings were allowed to proceed for 90 min on yeast extract-peptone-dextrose (YEPD) plates. For each mating, >100 zygotes were scored (22).

Molecular techniques. DNA and RNA manipulations, including plasmid DNA isolation and Northern blot analysis, were performed as described previously (43). Total RNA for microarray experiments and Northern blots was extracted from yeast cultures by using published procedures (44). Specific DNA fragments for Northern blot probes were prepared by PCR.

Plasmid construction was done by PCR-based in vivo recombination (37) and verified by DNA sequencing (Princeton Syn/Seq facility). The upstream *KAR3* minimal sequence was recombined into the expression vector pLG670-Z (15) using the forward primer 5'-GGA AAG CAG GAA AGG AAA AAA TTT TTA

GGC TCG AGC AAA ATG AAT CTC AAA CAA AAT CAA AAC AAA CAC-3' and the reverse primer 5'-ACG CTA TAT ATA CAC GCC TGG CGG ATC TGC TCG AGG TGT TTG TTT TGA TTT TGT TTG AGA TTC ATT TTG-3'. Mutations in the putative Kar4 consensus sites were introduced into the same expression vector using the following primers: k3minmut31 forward, 5'-GGA AAG CAG GAA AGG AAA AAA TTT TTA GGC TCG AGC AAA ATG AAT CTC AAA CAA AAT CAG AAC AAA CAC-3'; k3minmut31reverse, 5'-ACG CTA TAT ATA CAC GCC TGG CGG ATC TGC TCG AGG TGT TTG TTC TGA TTT TGT TTG AGA TTC ATT TTG-3'; k3minmut41 forward, 5'-GGA AAG CAG GAA AGG AAA AAA TTT TTA GGC TCG AGC AAA ATG AAT CTC AAA CAG AAT CAA AAC AAA CAC-3', and k3minmut41 reverse, 5'-ACG CTA TAT ATA CAC GCC TGG CGG ATC TGC TCG AGG TGT TTG TTT TGA TTC TGT TTG AGA TTC ATT TTG-3'.

The pGBD-*KAR4* and pGAD-*KAR4* constructs contain the *ADHI* promoter for the constitutive expression of in-frame fusions between *KAR4* and the GAL4 DNA-binding domain (GBD) or the GAL4 transcription activation domain (GAD). Constructs were made by PCR-based in vivo recombination using genomic DNA (37). The PCR primers were *KAR4*-FOR1 (AAAGAGATCGA ATTCCCGGGGATCCATATGGCATTCCAAGATCCAAC) and *KAR4*-REV1 (TACTACGATTCATAGATCTCTGCAGGTTTATTGTGCTTTTGT ACTGGACTTT). Vectors pGBD-C2 and pGAD-C2 (19) were linearized with BamHI and PstI and introduced into yeast with PCR-amplified *KAR4* DNA (6). Constructs were confirmed by restriction mapping and nucleotide sequencing (Princeton Syn/Seq facility). pGBD-C2 and pGBD-*KAR4* were transformed into PJ69-4A, a *MATa* yeast two-hybrid reporter strain (19), and an isogenic *ste12Δ* strain (MY10133), created by using pSUL16 (10). Pheromone-dependent expression was assayed by spotting fivefold serial dilutions of transformants, pregrown to exponential phase in liquid SC-TRP, onto SC-TRP and SC-TRP,-HIS agar plates, with or without α-factor at 3 μM. Plates were incubated for 2 to 5 days at 30°C. pGAD-C2 and pGAD-*KAR4* were transformed into S288C-derived isogenic strains MY3375 (*KAR4 STE12*) and MY4166 (*kar4Δ STE12*), as well as W303-derived isogenic strains MY3265 (*KAR4 STE12*) and MY5100 (*KAR4 ste12Δ*). All strains harbored a reporter construct with the *KAR3* minimal promoter region upstream of *lacZ* (pMR5620). β-Galactosidase assays on the minimal and mutated upstream region of *KAR3* were performed as previously described (23, 42).

Microarray and cluster analysis. Strains MS3213 (wild type) and MS3215 (*kar4Δ*) were grown to mid-exponential phase in YEPD (pH 3.5), after which α-factor or methanol (mock treatment) was added. The cultures were incubated for 90 min to allow full expression of Kar4 and other late genes involved in mating. RNA was extracted as described above, and the preparation of biotinylated cRNA, hybridization, and microarray scanning were performed according to the manufacturer's instructions (Affymetrix, Santa Clara, CA). Hybridization to the yeast genome S98 array was done in a GeneChip hybridization oven 640 at 45°C for 16 h, washing steps were performed by using the Affymetrix GeneChip fluidics station 400, and the GeneChips were scanned by using an Agilent GeneArray scanner. Raw data were normalized and clustered using the *k*-means clustering algorithm to group genes based on the similarity of their normalized gene expression profiles (49). The fold induction or repression for each of the obtained clusters was determined from the ratio of the raw data for each of the genes in the cluster (wild type to *kar4Δ* mutant).

To identify upstream Kar4 binding sites, the AlignACE algorithm was used (18, 48). The statistical significance of the sequence motif obtained is expressed as the maximum a priori log likelihood (MAP) score and the specificity score (18). The MAP score measures the degree to which a motif is over-represented relative to the expectation for the random occurrence of such a motif in the sequence under consideration. The specificity score is a measure of the degree to which the distribution of sites is skewed toward the input set, and it takes into account the sequence of all of the 5' upstream regulatory regions in the genome, highlighting motifs that are found preferentially in association with the genes under consideration (18).

Temporal regulation of Kar4-dependent genes. The 29 Kar4-dependent and 34 Kar4-independent genes showing >2.5-fold induction by pheromone were selected for analysis. The data for the timing of induction in a *bar1Δ* strain for these genes (0, 15, 30, 45, and 60 min after induction) by 50 nM pheromone were obtained from previously published data (39). Similarly, data for the dose response of a *bar1Δ* strain to alpha factor exposure (0.15, 0.5, 1.5, 5, 15.8, 50, and 158 nM pheromone) after 30 min were previously published (39). A Stineman function was fit to all datum points to obtain a local smoothing (KaleidaGraph 4.01; Synergy Software). The datum points corresponding to the half-maximal value (*t*_{1/2}) were interpolated, and the *t*_{1/2} for each gene was obtained. The statistical significance was tested by using a heteroscedastic Student *t* test.

TABLE 1. Strains and plasmids used in the present study^a

Strain or plasmid ^a	Genotype or relevant markers ^b	Source or reference
Strains		
MS3212	<i>MATα leu2-3,112 his3Δ200 ura3-52 ade2-101 kar4Δ::HIS3</i>	
MS3213	<i>MATα trp1Δ1 his3Δ200 ura3-52 ade2-101</i>	
MS3215	<i>MATα leu2-3,112 his3Δ200 ura3-52 ade2-101 kar4Δ::HIS3</i>	
MS3216	<i>MATα leu2-3,112 his3Δ200 ura3-52 ade2-101 kar4Δ::HIS3</i>	
MY3265	<i>MATα ura3-52 trp1 his4-519 leu2 can1-101</i>	
MY3375	<i>MATα ura3-52 leu2Δ1 his3Δ200</i>	
MY4166	<i>MATα kar4Δ ura3-52 leu2Δ1 his3Δ200</i>	
MY5100	<i>MATα ste12Δ::leu2Δ1 ura3-52 trp1 his4-519 leu2 can1-101</i>	
MY5792	<i>MATα ura3-52 leu2Δ1 his3Δ200 KAR4::HA</i>	
MY10133	<i>MATα ste12Δ::LEU2 trp1-901 leu2-3,112 ura3-52 his3Δ200 gal4Δ gal80Δ GAL2-ADE2 LYS2::GAL1-HIS3 met2::GAL7-lacZ</i>	
PJ69-4A	<i>MATα trp1-901 leu2-3,112 ura3-52 his3Δ200 gal4Δ gal80Δ GAL2-ADE2 LYS2::GAL1-HIS3 met2::GAL7-lacZ</i>	19
BY4741	<i>MATα his3Δ1 leu2Δ0 met15Δ0 ura3Δ0</i>	49
BY4742	<i>MATα his3Δ1 leu2Δ0 lys2Δ0 ura3Δ0</i>	49
Y01430	<i>MATα prm2Δ::KanMX his3Δ1 leu2Δ0 met15Δ0 ura3Δ0</i>	49
Y11430	<i>MATα prm2Δ::KanMX his3Δ1 leu2Δ0 lys2Δ0 ura3Δ0</i>	49
Y03462	<i>MATα kar4Δ::KanMX his3Δ1 leu2Δ0 met15Δ0 ura3Δ0</i>	49
Y13462	<i>MATα kar4Δ::KanMX his3Δ1 leu2Δ0 lys2Δ0 ura3Δ0</i>	49
Plasmids		
pLG669-Z	<i>lacZ 2μm URA3</i>	15
pMR2806	<i>P_{KAR3-L11}-lacZ 2μm URA3</i>	
pMR2802	<i>P_{KAR3-L16}-lacZ 2μm URA3</i>	
pMR2892	<i>P_{KAR3-L18}-lacZ 2μm URA3</i>	
pMR2808	<i>P_{KAR3-L14}-lacZ 2μm URA3</i>	
pMR2896	<i>P_{KAR3-L15}-lacZ 2μm URA3</i>	
pMR5620	<i>P_{KAR3-min}-lacZ 2μm URA3</i>	
pMR5621	<i>P_{KAR3-min31-2}-lacZ 2μm URA3</i>	
pMR5622	<i>P_{KAR3-min41-2}-lacZ 2μm URA3</i>	
pMR2654	<i>KAR4::HA CEN ARS URA3</i>	
pMR2516	<i>KAR4 CEN ARS URA3</i>	
pRS416	<i>CEN ARS URA3</i>	45
pMR5511	<i>kar4^{C214GC217C}::HA CEN ARS URA3</i>	
pMR3356	<i>P_{GAL}-KAR4::HA-short CEN ARS URA3</i>	
pMR3459	<i>P_{GAL}-KAR4::HA-long CEN ARS URA3</i>	
pGBD-C2	<i>GBD TRP1 2μm</i>	19
pMR4997	<i>GBD-KAR4 TRP1 2μm</i>	
pGAD-C2	<i>GAD LEU2 2μm</i>	19
pMR5000	<i>GAD-KAR4 LEU2 2μm</i>	
pMR5665	<i>P_{ter}-6xHN-KAR4; Cm^r</i>	
pMR5669	<i>P_{ter}-6xHN-STE12; Cm^r</i>	
pPROTet.E133	<i>P_{ter}-6xHN; Cm^r</i>	Clontech
pSUL16	<i>ste12Δ::LEU2</i>	11

^a Unless stated otherwise, all strains and plasmids were constructed in this laboratory.

^b Cm^r, chloramphenicol resistant.

Detection of Kar4. Hemagglutinin (HA) epitope-tagged forms of Kar4 (14, 23) were detected with 12CA5 monoclonal antibody (Princeton monoclonal facility). Total yeast protein was extracted from cells as previously described (36). Proteins were run on a 10% polyacrylamide gel at 100 V and transferred to supported nitrocellulose membrane (Optitran; pore size, 0.2 μ m; Schleicher & Schuell) at 100 V for 60 min. Western blotting was performed as previously described (14), and the Kar4::HA proteins were detected by using the ECL kit (Amersham, Arlington Heights, IL).

Chromatin immunoprecipitation assay. Strain MY5792, which contains a chromosomal version of *KAR4* with the HA epitope inserted at the carboxyl terminus of the coding region (14), and an untagged control strain MY3375, were grown to an optical density at 600 nm (OD₆₀₀) of 0.7. Cells were treated with α -factor dissolved in methanol or methanol alone for 90 min. Chromatin immunoprecipitation was performed as described previously (21), except that formaldehyde cross-linking was carried out for 1 h at room temperature. The primers used to amplify the upstream region of *KAR3* were 5'-TTC GGC TTC TCT ATC GCT TT-3' and 5'-TGC CAG AAT TTC TCT AAG TCC-3'.

Gel shift assay. The open reading frames of *KAR4* and *STE12* were amplified from genomic DNA by PCR (6) and inserted into pPROTet.E133 vector (Clontech, Palo Alto, CA) to create in-frame fusions to a tag of six repeated His-Asn residues (6xHN). Fusion proteins were expressed in *Escherichia coli* (BL21Pro) by induction with 100 ng/ml of anhydrotetracycline (Clontech) for 4 h and partially purified by using NTA spin columns (QIAGEN, Valencia, CA). Cells containing the pPROTet.E133 vector were processed in parallel as a control for background DNA-binding activity. Protein concentrations were determined by using the Bradford protein assay (Bio-Rad Laboratories, Hercules, CA) with bovine serum albumin as the protein standard. Protein purity was assessed by using sodium dodecyl sulfate-polyacrylamide gel electrophoresis and Coomassie blue staining (1).

DNA gel shift assays were conducted according to the method of Dolan et al. (10) with minor modifications. DNA probes were prepared by annealing complementary oligonucleotides encoding the minimal Kar4-dependent region contained in the *KAR3* promoter plus five downstream nucleotides to include the full Kar4-responsive consensus sequence. The 5' overhangs were filled with Klenow

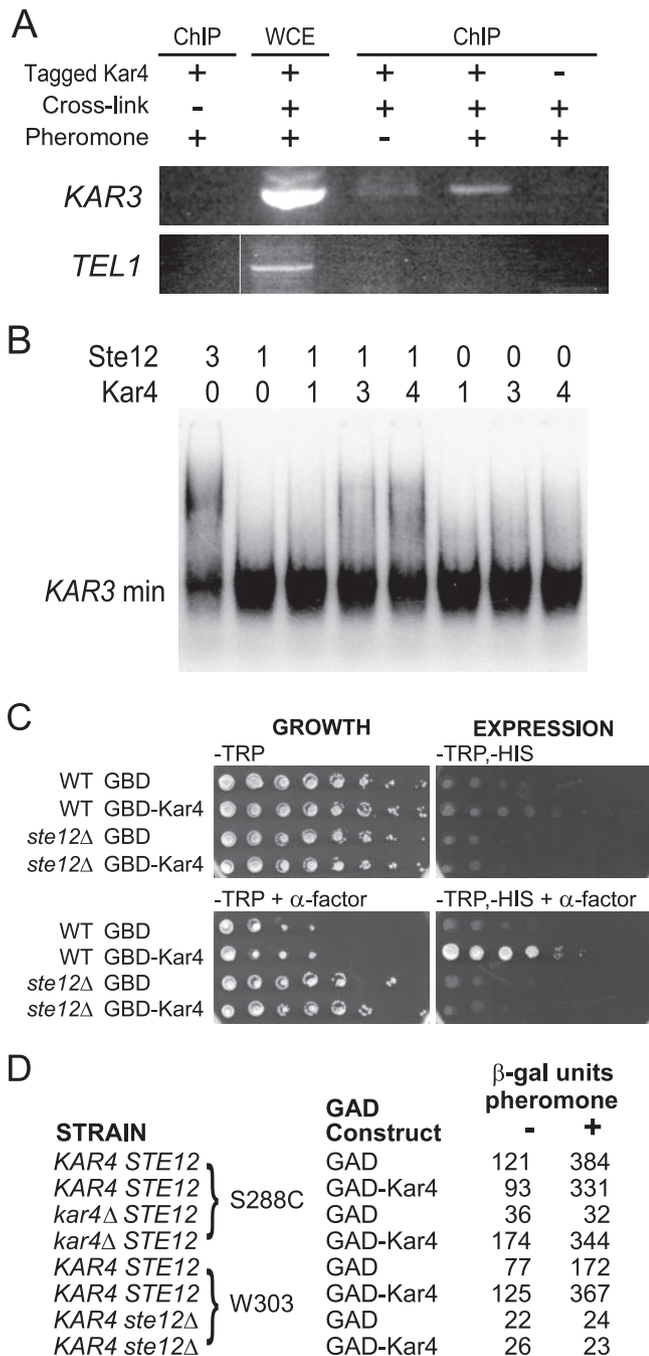


FIG. 1. (A) Kar4 functions at the promoter region of *KAR3*. Chromatin immunoprecipitation of Kar4::HA-bound upstream sequence of *KAR3*. Strain MY5792 harboring a chromosomal version of *KAR4::HA* (tagged Kar4) and the untagged control strain MY3375 were either treated with pheromone for 90 min or mock treated with MeOH. Immunoprecipitated (ChIP) or whole-cell extract (WCE) DNA was amplified with primers specific to upstream sequences of *KAR3* and *TEL1* (nonspecific control). PCR fragments were fractionated and visualized by using conventional agarose gel electrophoresis. Lane 1, MY5792 no-cross-link control; lane 2, MY5792 whole-extract control; lane 3, cross-linked tagged strain MY5792, not induced by pheromone; lane 4, pheromone-induced cross-linked MY5792; lane 5, cross-linked, pheromone-induced untagged strain MY3375. (B) Kar4 and Ste12 bind to the *KAR3* promoter. The results of a gel shift assay using radioactively labeled *KAR3* minimal promoter region (*KAR3-min*) in the presence of various amounts of *E. coli* expressed Ste12 and Kar4 is

fragment (New England Biolabs), dCTP, dTTP, dGTP, and ³²P-α-dATP (Amersham) according to the enzyme manufacturer's specification. For binding experiments, poly(dI-dC) competitor was added to a final concentration of 100 μM. Protein extracts were added to a constant final concentration in each experiment (typically between 3 and 6 μg per reaction). Where Ste12 or Kar4 concentrations were varied, compensatory amounts of equivalent protein from cells bearing the vector plasmid were added.

Site-directed mutagenesis. Site-directed mutagenesis of the zinc-finger domain in the 3X HA-tagged version of *KAR4* (pMR2654) was performed by using the QuikChange II XL kit (Stratagene, La Jolla, CA). The primers used to mutate the putative zinc-finger domain of *KAR4* were 5'-AAA AAT GCA ATG GCA CGG TTG GAT GGG TAT CAC AGG TAC A-3' (sense) and 5'-TGT ACC TGT GAT ACC CAT CCA ACC GTG CCA TTG CAT TTT T-3' (anti-sense). The pMR5511 construct was sequenced to confirm the change (Princeton Syn/Seq facility).

Galactose induction of Kar4. For galactose induction experiments, the *kar4Δ* strain MS3216 harboring the plasmids pMR3356 (*P_{GAL}-KAR4-short::HA*), pMR3459 (*P_{GAL}-KAR4-long::HA*), and pMR2654 (*KAR4::HA*) were grown in synthetic complete media without uracil, with 2% raffinose, to mid-exponential phase. Cultures were induced by the addition of 2% galactose plus 0.35% glucose (for *KAR4-long::HA*) and 2% galactose plus 0.2% glucose (for *KAR4-short::HA*). Galactose induction was for 3 h, after which pheromone was added. Aliquots of cells were harvested at each time point (0, 5, 10, 20, 30, 45, 60, 75, and 90 min after addition of pheromone) and snap-frozen in liquid nitrogen. Northern blotting was performed as previously described.

RESULTS

Kar4 binds to DNA upstream of the *KAR3* gene. Although Kar4 was implicated as a transcriptional activator of *KAR3* and *CIK1*, it was not determined whether Kar4 binds to DNA along with Ste12 or acts indirectly to facilitate Ste12's association or activity. As a first test, we used chromatin immunoprecipitation to detect the association of epitope-tagged Kar4 with the upstream regulatory region of *KAR3*. Strains expressing Kar4::Hap or untagged Kar4 were treated with α-factor or mock treated with methanol, the DNA was cross-linked, and the chromatin containing Kar4 was immunoprecipitated (Fig. 1A). Enrichment of *KAR3*-specific DNA sequences was detected by PCR and compared to a nonspecific DNA sequence from the telomere of chromosome 1 (*TEL1*).

A small amount of *KAR3* upstream DNA, but not *TEL1*, was precipitated from vegetative cells. The addition of pheromone caused a significant enrichment of the *KAR3* DNA above the level observed in vegetative cells, a finding consistent with increased levels of Kar4. Immunoprecipitation by the HA an-

shown. The relative amount of each protein is indicated. (C) Kar4 binding confers pheromone inducible expression. The wild-type yeast two-hybrid reporter strain PJ69-4A (WT) or an isogenic strain containing a *ste12Δ* were transformed with either the GBD vector plasmid or with GBD-Kar4. Strains were grown to exponential phase, and serial dilutions were spotted onto selective plates with or without 3 μM pheromone (α-factor) dissolved in MeOH. All cells can grow on -TRP plates (GROWTH). Only cells expressing the *GAL-HIS3* reporter gene can grow on -TRP,-HIS plates (EXPRESSION). (D) Fusing an activation domain to Kar4 does not remove the requirement for Ste12. S288C-derived strains MY3375 (*KAR4 STE12*) and MY4166 (*kar4Δ STE12*), as well as W303-derived strains MY3265 (*KAR4 STE12*) and MY5100 (*KAR4 ste12Δ*), were transformed with either the GAD vector plasmid or with the GAD-Kar4 plasmid. Each strain also harbored the reporter construct with the *KAR3* minimal promoter region fused upstream of *lacZ* (pMR5620). Expression from the promoter is presented in β-galactosidase (β-gal) units from cultures grown in the presence (+) or absence (-) of 6 μM α-factor (pheromone).

as that caused by complete deletion of *KAR4*, this domain is clearly important for Kar4 function.

Kar4 regulates the expression of many genes. We performed microarray experiments to determine whether Kar4 has a more global effect on the transcriptional regulation of genes involved in the yeast pheromone response. cRNA prepared from total RNA of pheromone or mock-treated wild-type and *kar4Δ* cells and used to probe Affymatrix yeast microarrays in duplicate. Raw hybridization data were collected, normalized, and clustered into 20 original clusters according to a *k*-means clustering algorithm, grouping the genes based on the similarity of their normalized gene expression profiles (48). Clusters containing genes that showed neither pheromone-dependent regulation nor evidence of Kar4-dependent expression were not examined further. Clusters with similar patterns of expression were pooled together into seven new clusters (Fig. 3). Clusters 1 (414 genes) and 2 (850 genes) are comprised of pheromone-regulated genes whose induction or repression was independent of Kar4. The remaining clusters were filtered by the ratio of gene expression in the *kar4Δ* mutant relative to the wild type; only genes showing a >2-fold difference were further analyzed. Cluster 3 contains 70 genes whose expression was reduced in the *kar4Δ* mutant after pheromone treatment. Two expected genes within this cluster are *KAR3* and *CIK1*. In addition to these three anticipated gene clusters, four other clusters were unexpected. Cluster 4 contains 225 genes that were expressed more strongly after pheromone induction in the *kar4Δ* mutant, indicating a possible role for Kar4 in the repression of genes after exposure to pheromone. Two clusters contained genes whose expression is constitutive in wild-type cells but which are either repressed (cluster 5) or induced (cluster 6) by pheromone in the *kar4Δ* mutant. These clusters suggest that Kar4 plays a specific role in maintaining the expression of these genes at normal levels during the pheromone response. Cluster 7 contained 55 genes whose vegetative expression was elevated in the *kar4Δ* mutant but which showed constitutive expression in wild-type cells, a finding consistent with Kar4 having a significant regulatory function during vegetative growth.

To determine whether the clustering of genes based on their pattern of Kar4 dependence reflected a deeper functional relationship, we examined the gene annotations for significant correlations by using FunSpec (40). Cluster 3 was significantly enriched in genes previously shown to be involved in nuclear congression (e.g., *KAR3* and *CIK1*) or known to affect microtubule function (e.g., *SPC25*) or else implicated in the structure of the cytoskeleton (e.g., *YOR129C*), a finding consistent with Kar4's function in regulating genes involved in nuclear congression ($P < 0.0004$ [FunSpec]). Other genes within the pheromone-induced, Kar4-dependent cluster may have roles in mating that are not directly related to microtubule function, such as *PRM2*, which encodes a putative pheromone-induced membrane protein (17). Although no function has been previously reported for *PRM2*, we found that karyogamy failed in 30% of *prm2Δ/prm2Δ* zygotes, a finding consistent with a role in mating (Fig. 4A). Many of the remaining genes in the Kar4-dependent cluster are not characterized. Kar4 most likely regulates genes in which both mating partners must be mutant to express the mating defect, as is the case for both *KAR3* and *CIK1*. Such "bilateral" mutants and mutants with less severe

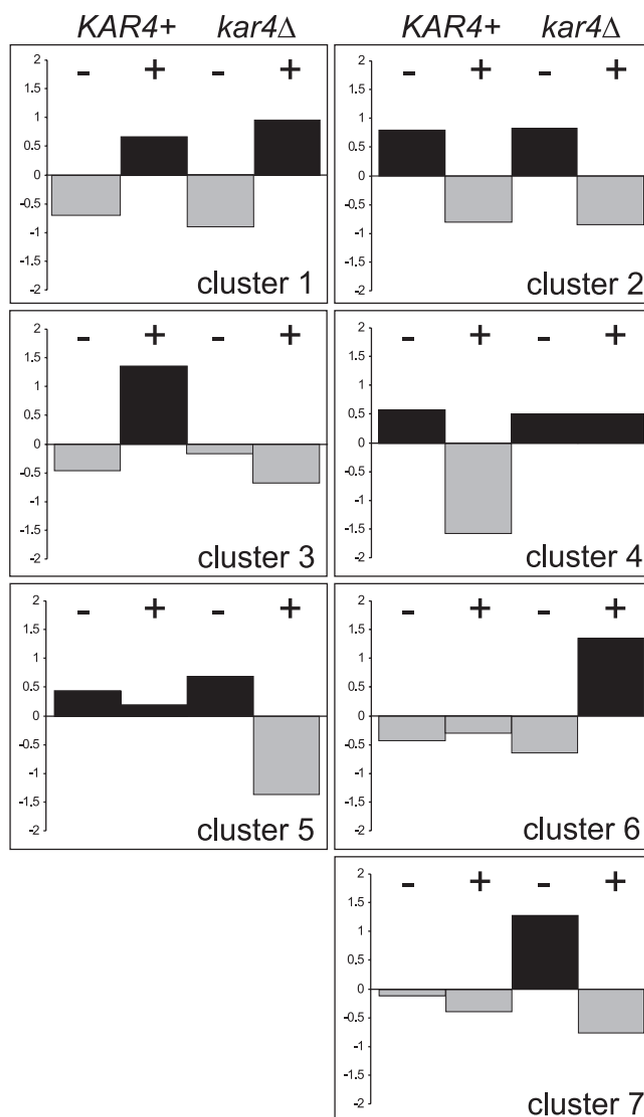


FIG. 3. Kar4-dependent clusters after pheromone treatment. Strains MS3213 (wild type) and MS3215 (*kar4Δ*) were exposed to pheromone (+) or mock treated with methanol (-) for 90 min. RNA was extracted and prepared for microarray analysis as described in Materials and Methods. Genes were clustered according to their normalized expression profiles. Specifically, we identified seven clusters that show Kar4-independent pheromone induction (cluster 1), Kar4-independent repression in pheromone (cluster 2), Kar4-dependent pheromone induction (cluster 3), Kar4-dependent repression in pheromone (cluster 4), constitutive expression in wild-type cells but repression by pheromone in the *kar4Δ* strain (cluster 5), constitutive expression in wild-type cells but induction by pheromone in the *kar4Δ* strain (cluster 6), and constitutive expression in wild-type cells but higher levels of mitotic expression in the *kar4Δ* strain (cluster 7).

defects would have been missed in previous screens for mutations affecting mating (22, 41).

Cluster 4 contained genes whose expression was reduced during mating in a Kar4-dependent manner. Cluster 4 was enriched in genes involved in cell wall functions (e.g., *CIS3/PIR2*, *CWP1*, *SKG1*, *MHP1*, and *YDL222C*) or which act as molecular chaperones (e.g., *SSE2*, *HSP30*, *HSP42*, *SIS1*, *YNL077W*, *HSP78*, *SSA4*, *HSP104*, and *HSP82*). Downregula-

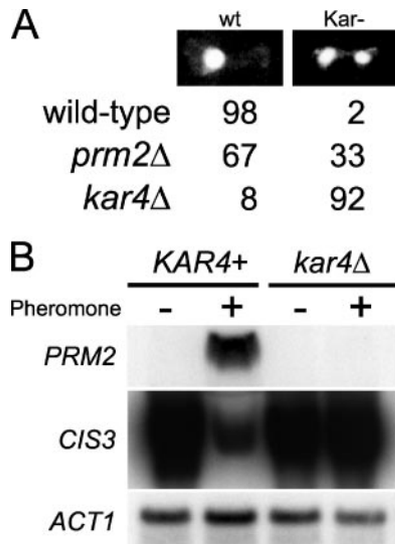


FIG. 4. Kar4-dependent induction and repression of genes. (A) *PRM2* is required for efficient nuclear fusion. Three different mating mixtures (wild-type \times wild-type, $n = 55$; *kar4*Δ \times *kar4*Δ, $n = 148$; and *prm2*Δ \times *prm2*Δ, $n = 195$) were examined for karyogamy defects. Mating mixtures were stained with DAPI, and the phenotypes were observed and scored as fused (wt) or unfused (*Kar*⁻) nuclei. The percentages of each class are shown. (B) Northern blot analysis of wild type (MS3213) and *kar4*Δ (MS3215) mRNA in the absence (-) or presence (+) of pheromone. The blot was hybridized to a probe for *PRM2* (upper panel) or *CIS3* (middle panel). A duplicate Northern blot was hybridized with an *ACT1* probe (bottom panel).

tion of cell wall proteins would be consistent with the requisite removal of cell wall at the zone of cell fusion during mating (13).

To confirm the microarray data, Northern blots were performed on two representative genes from clusters 3 and 4 (Fig. 4B). *PRM2* was observed to be strongly induced by pheromone, in a *Kar4*-dependent way. Similarly, *CIS3* was strongly repressed by the addition of pheromone, but levels remained high in the *kar4*Δ mutant.

Identification of a consensus *Kar4* regulatory motif. To determine whether the *KAR3*-min region is both necessary and sufficient for *Kar4*-mediated pheromone induction of *KAR3*, the 30-bp segment was synthesized and placed upstream of a *lacZ* reporter gene (Fig. 5A). In the presence of pheromone, the 30-bp region drove the expression of the reporter gene to the same extent as the full-length *KAR3* promoter (L11) (23), indicating that this 30-bp minimal sequence contained all of the necessary regulatory information. It was previously noted that the 30 bp contains a repeated sequence motif (CAAA), which was suggested to be a regulatory element for *Kar4*, in addition to a weak *Ste12*-binding site (23).

Two *KAR3* promoter deletions, L14 (pMR2802) and L15 (pMR2896), with endpoints within one of the CAAA repeats, had previously been found to destroy pheromone induction, which is consistent with an essential role for this element in *Kar4*-dependent pheromone regulation (Fig. 5A). Mutation of one of the A residues within this CAAA repeat eliminated pheromone induction of the *lacZ* reporter gene (Fig. 5C). In contrast, mutation of an equivalent A, adjacent to the PRE

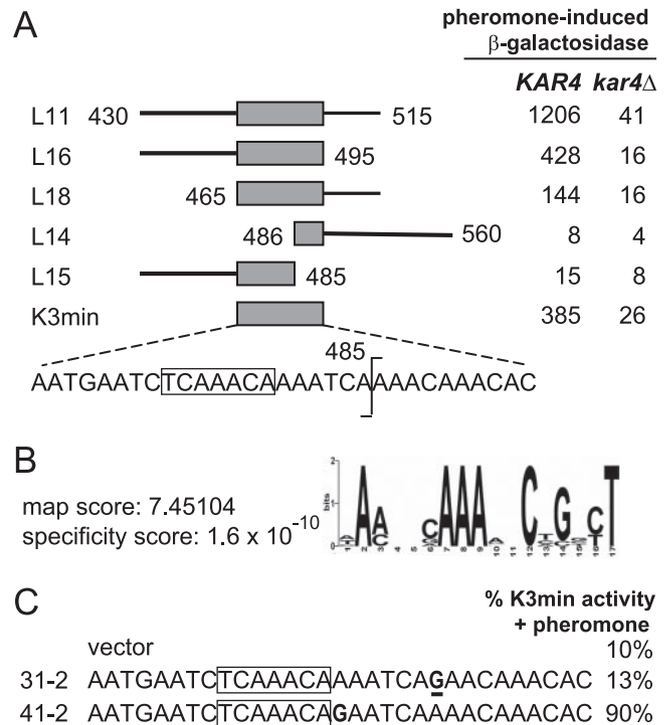


FIG. 5. Defining *Kar4* activation parameters. (A) *KAR4*-dependent minimal regulatory region. The DNA region upstream of the *lacZ* gene, lacking upstream activator sequences. Expression is presented as units of β -galactosidase specific activity in the presence of mating pheromone. Assays were performed with MY3265 (*KAR4*) or MY4166 (*kar4*Δ) harboring plasmids with deletions in the *KAR3* promoter regions. L11 (pMR2806), L16 (pMR2802) and L18 (pMR2892) delimit the *KAR4*-dependent region to a 30-bp region upstream of the *KAR3* gene. Flanking the lines are numbers that correspond to the endpoints of the deletions within the original 560-bp region (23). K3 min (pMR5620) contains the minimal 30-bp *KAR4*-dependent region indicated by the shaded box. The nucleotide sequence is shown below K3 min. The vertical line at position 485 indicates the endpoints for deletions L14 (pMR2808) and L15 (pMR2896). The results for L11, L14, L15 L16, and L18 were published previously (23). (B) Sequence alignment of upstream DNA from *Kar4*-regulated genes as identified by AlignAce analysis. Upstream regulatory regions of *Kar4*-dependent pheromone-induced genes showing a fourfold or greater induction were analyzed for sequence enrichment. See the Materials and Methods for descriptions of the MAP score and the specificity score. (C) Mutational analysis of the *Kar4*-dependent region. The results of β -galactosidase assays represented as a percentage of the *KAR3*-minimal region promoter activity (% K3 min) in the presence of mating pheromone are shown. MY3265 harboring plasmids containing mutated (31-2, pMR5621; 41-2, pMR5622) or wild-type (pMR5620) minimal *KAR3* upstream regulatory region, as well as the vector control (pLG669-Z, vector), were analyzed for promoter activity. Mutated nucleotides are underlined. The consensus pheromone response element (PRE) is boxed.

(TCAAACA), within an overlapping CAAA, had no effect on induction (Fig. 5C).

To determine whether the CAAA sequence is also upstream of other genes regulated by *Kar4*, we compared the DNA sequences of the 15 genes in cluster 3 whose expression after pheromone induction was reduced at least fourfold or more in the *kar4*Δ mutant relative to the wild type. These genes included *KAR3*, *CIK1*, *SPC25*, *PRM2*, *ISC1*, *IME4*, *CDA4*, *JJJ2*, *ECM23*, *YPL033C*, *YDR281C*, *YOR129C*, *YNL033W*, *YJL218W*,

TABLE 2. Kar4-dependent and -independent genes induced ≥ 2.5 -fold based on microarray data

Gene type	No. of genes	Genes
Kar4-dependent genes	29	<i>KAR3</i> , <i>CIK1</i> , <i>PRM2</i> , <i>IME4</i> , <i>SPC25</i> , <i>YOR129C</i> , <i>YNL105W</i> , <i>YPL033C</i> , <i>YIL218W</i> , <i>ECM23</i> , <i>JJJ2</i> , <i>YNL033W</i> , <i>CDA2</i> , <i>ISC1</i> , <i>FUI1</i> , <i>MMP1</i> , <i>PCL2/CLN4</i> , <i>RPO31</i> , <i>MHT1</i> , <i>HXT9</i> , <i>CLN3</i> , <i>YGL204C</i> , <i>PGM1</i> , <i>YDR340W</i> , <i>SCW11</i> , <i>CUP9</i> , <i>RCL1</i> , <i>HOS2</i> , <i>YOR225W</i>
Kar4-independent genes	34	<i>PRM6</i> , <i>FUS2</i> , <i>YDR124W</i> , <i>PRM1</i> , <i>ASG7</i> , <i>YOR343C</i> , <i>FIG1</i> , <i>YIL080W</i> , <i>FIG2</i> , <i>FUS1</i> , <i>KAR5</i> , <i>RRN11</i> , <i>PRM5</i> , <i>SST2</i> , <i>HYM1</i> , <i>DIG2</i> , <i>FAR1</i> , <i>YJL169W</i> , <i>SUT1</i> , <i>HMI1</i> , <i>YNL324W</i> , <i>BOS1</i> , <i>BOP3</i> , <i>YML048W-A</i> , <i>ORT1</i> , <i>ECM13</i> , <i>AGA1</i> , <i>MFA1</i> , <i>GPA1</i> , <i>KTR2</i> , <i>PRP39</i> , <i>YMR304C-A</i> , <i>AGA2</i> , <i>STE2</i>

and *YNL105W*. Using AlignACE (18) to identify DNA sequence motifs that are overrepresented upstream of the input genes, we found significant enrichment of one motif, C/TAA NNCAAANNCGNYT (Fig. 5B, MAP score of 7.45104 and specificity score of 1.6×10^{-10}). This motif was found upstream of 9 of the input genes, including *KAR3* (CAAAACA AACACTGTTT). As discussed above, mutation of the second A of the conserved motif destroyed promoter activity from the *KAR3* minimal activating sequence (Fig. 5C). Two other putative motifs encompassed the same CAAA sequence but exhibited low specificity scores due to the high TA content (motif 1, MAP score of 28.9849, specificity score of 7.2×10^{-1} ; motif 2, MAP score of 18.691, specificity score of 1.1×10^{-1}).

Temporal regulation by Kar4. The additional requirement of Kar4 along with Ste12 for the regulation of genes during mating provides a potential mechanism for temporal regulation during the pheromone response. From the microarray data, we generated lists of the genes that were induced by pheromone at least 2.5-fold in both the Kar4-independent cluster and the Kar4-dependent cluster (Table 2). To compare the two sets of genes, published gene expression profiles were compared with respect to both the kinetics of induction and the dose response to pheromone (39). For each gene we determined the time required for half-maximal induction ($t_{1/2}$). For the set of 34 Kar4-independent genes, ca. 25% showed half-maximal inductions of 3 min or less after addition of pheromone and ca. 75% of genes showed $t_{1/2}$'s of 9 min or less, with a median $t_{1/2}$ of 7.5 min (Fig. 6A). The rapid induction of this class of genes is consistent with previous data showing that the prototype pheromone-induced gene, *FUS1*, does not require protein synthesis for its induction (32). In contrast, the set of 29 Kar4-dependent genes were delayed in their response to pheromone. Only 25% showed half-maximal induction by 10 min after pheromone addition and 75% required as long as 16 min, with a median $t_{1/2}$ of 12 min. Thus, the requirement for Kar4 imposed a significant delay in pheromone induction to this class of genes ($P = 3.3 \times 10^{-7}$).

The two sets of genes also showed significant differences with respect to the pheromone dose response. In the *bar1Δ* strain used in the study, Kar4-independent genes required a median concentration of only 1.25 nM pheromone to achieve half-maximal expression (Fig. 6B). In contrast, the Kar4-dependent genes required a significantly higher median concentration of pheromone (5.2 nM) to reach half-maximal expression ($P = 1.59 \times 10^{-5}$).

These observations suggest that Kar4 specifically imparts a delayed response to genes regulated by mating pheromone. To test this hypothesis, we examined the effect of precocious expression of Kar4 on the temporal regulation of genes that are either induced (*KAR3* and *PRM2*) or repressed (*CIS3*) by pheromone, depending on Kar4. To perturb the kinetics of induction, Kar4 was exogenously regulated using the galactose-inducible *GAL1* promoter. We used constructs expressing one

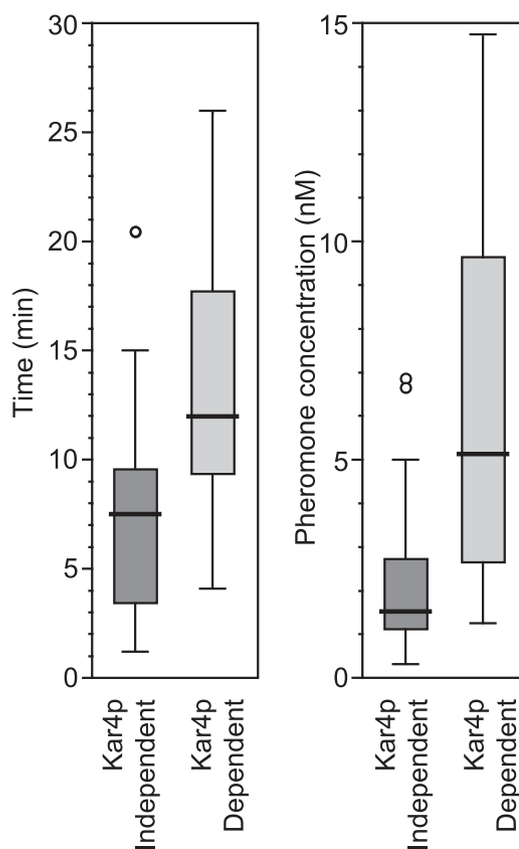


FIG. 6. Temporal and pheromone concentration regulation of Kar4-dependent genes. (A) Genes showing greater than 2.5-fold induction by pheromone from cluster 1 (Kar4-independent [34]) and cluster 2 (Kar4 dependent [29]) were selected, and datum points were taken representing the \log_2 (ratio) of normalized expression at 0, 15, 30, 45, and 60 min after induction by pheromone (39). A pheromone response curve was generated for each gene, and the $t_{1/2}$ was extracted from the interpolated data. Primary data were from Roberts et al. (39). (B) Box plot representing the concentration of pheromone corresponding to the half-maximal induction of each of the Kar4-dependent and Kar4-independent pheromone response genes. The data determined from the \log_2 (ratio) of normalized expression at 30 min after exposure of a *bar1Δ* strain to various concentrations of pheromone (0.15, 0.5, 1.5, 5, 15.8, 50, and 158 nM α -factor) Primary data were from Roberts et al. (39).

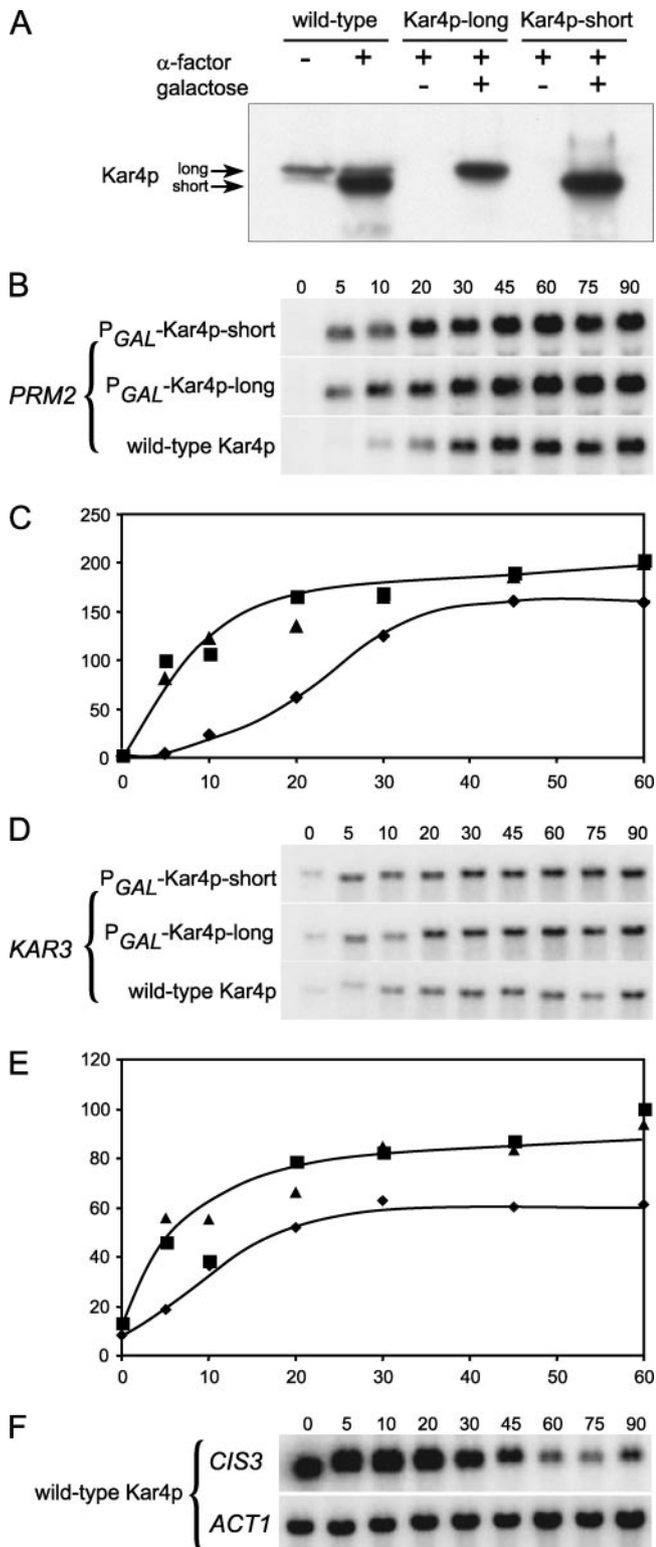


FIG. 7. Kar4-dependent temporal regulation of *PRM2*, *KAR3*, and *CIS3*. (A) Western blot comparison of pheromone- and galactose-induced Kar4::HA. The MS3216 *kar4Δ::HIS3* strain harbored either the wild-type *KAR4::HA* (pMR2654), *P_{GAL}-KAR4-short::HA* (pMR3356), or *P_{GAL}-KAR4-long::HA* (pMR3459) plasmid constructs. The cells were grown in the absence (–) or presence (+) of galactose and in the absence (–) or presence (+) of pheromone (α-factor). The arrows indicate the two forms of Kar4::HA. (B to F) Time course of

of the two forms of Kar4 from the *GAL1* promoter (*P_{GAL}*). The shorter, pheromone-induced form was expressed from *P_{GAL}-KAR4-short::HA*, and the longer, constitutively expressed form was expressed from *P_{GAL}-KAR4-long::HA*. In both cases the concentrations of glucose and galactose were adjusted to achieve levels of the *P_{GAL}*-expressed proteins comparable to those observed for the endogenous short form after pheromone induction (14). Protein levels of the endogenous (wild-type) and galactose-induced forms of Kar4 were verified to be similar (Fig. 7A). All strains were grown in galactose and then exposed to mating pheromone. At various time intervals, RNA was extracted, blotted, and probed for expression of *PRM2* (Fig. 7B and C), *KAR3* (Fig. 7D and E), and *CIS3* (Fig. 7F).

In wild-type cells, *PRM2* mRNA levels were not detected prior to the pheromone treatment. After pheromone addition, *PRM2* mRNA levels increased gradually, reaching half-maximal levels after approximately 25 min (Fig. 7B and C). In both strains in which Kar4 was expressed from *P_{GAL}*, initial mRNA levels were not detected but increased very rapidly after pheromone addition, reaching half-maximal levels within 5 min (Fig. 7B and C). Plateau levels of *PRM2* were somewhat elevated relative to the wild type.

In wild-type cells, a basal mitotic level of *KAR3* mRNA was observed prior to pheromone addition (Fig. 7D and E). The *KAR3* mRNA levels increased more quickly than *PRM2*, reaching half-maximal levels approximately 10 min after pheromone addition. In both strains in which Kar4 was expressed from *P_{GAL}*, the initial levels of *KAR3* mRNA were similar to that of the wild type but increased more rapidly, reaching half maximal levels with 5 min after pheromone addition (Fig. 7D and E). In addition, the plateau levels of *KAR3* mRNA were significantly elevated compared to cells with endogenously expressed Kar4. Thus, the kinetics of both *PRM2* and *KAR3* induction were significantly accelerated by the premature expression of Kar4, albeit to different extents.

Either form of Kar4 when prematurely produced was able similarly to induce and alter the kinetics of *PRM2* and *KAR3* expression. These observations confirm that the two proteins are functionally equivalent but differ mainly in their patterns of expression and stability (14).

In contrast to *KAR3* and *PRM2*, *CIS3* mRNA levels were high in mitotic cells and began to decline after 20 to 30 min after pheromone addition, reaching levels three- to fourfold lower (Fig. 7F). Premature expression of Kar4 from *P_{GAL}* had no significant effect on the rate of mRNA decline (not shown), suggesting that the half-life of the mRNA is longer than any potential shift in the timing of repression. *ACT1* mRNA levels were essentially constant throughout the time of the experi-

kar4Δ cells (MS3216) harboring plasmid pMR2654 (◆, wild-type Kar4) or the galactose-induced plasmids pMR3356 (■, galactose-induced short form of Kar4, *P_{GAL}-Kar4-short*) or pMR3459 (▲, galactose-induced long form of Kar4 predominant, *P_{GAL}-Kar4-long*). Cells were grown in medium containing galactose, and pheromone was added at *t* = 0. Cells were harvested at the indicated times, and RNA was prepared. Northern blots were hybridized with probes to either *PRM2* (B), *KAR3* (D), *CIS3* (F), or *ACT1* (F). The graphs in panels C and E show the levels of mRNA in arbitrary intensity units.

ment, indicating that equal concentrations of RNA were analyzed at each time point (Fig. 7F).

DISCUSSION

Originally identified as a regulator of two specific genes required for karyogamy, we find that Kar4 has a more widespread role in the yeast mating response. In addition to being required for the pheromone-dependent induction of *KAR3* and *CIK1*, Kar4 is required for both the positive and the negative regulation of a number of genes during both mating and mitosis (Fig. 8A).

Among the genes positively regulated by Kar4 during mating is *PRM2*, encoding a member of the pheromone-regulated putative integral membrane proteins (PRM). We have found that Prm2 is required for efficient karyogamy, and it is likely that other proteins in this cluster will also have roles in mating.

An unexpected cluster contained genes repressed by pheromone in a Kar4-dependent manner. The observation that a gene is repressed by pheromone may reflect either a specific regulatory response to pheromone or downregulation of the gene in the G₁ phase of the cell cycle. However, because the *kar4Δ* strain arrests in G₁ like the wild-type strain, genes dependent on Kar4 for pheromone-dependent downregulation are showing a specific response to pheromone. Many genes in this cluster encode proteins that are either predicted to be membrane localized or are involved in cell wall function, including *CIS3/PIR2*, *HOC1*, *CWPI*, *YBR209W*, *CRC1*, *PHO5*, *PDR15*, *YGL146C*, *IZH4*, *YOL013W-A*, *FMP45*, *HBT1*, *RSN1*, *YSR3*, *HXT16*, *EDE1*, *ECM30*, *OPT1*, *MAL11*, *ECI1*, *EMP70*, *PST1*, *TGL1*, *SIM1*, *JSN1*, *LST8*, *YKL008C*, *SNX4*, *STD1*, *SKG1*, and *YNL181W*. One interpretation of this cluster of genes is that cell wall biosynthesis may be negatively regulated at the shmoo tip to allow efficient cell fusion during mating. Consistent with this hypothesis, the shmoo tip has reduced levels of β-1-3 glucan that are correlated with the efficiency of mating (13).

The cluster of genes exhibiting Kar4-dependent pheromone repression was also enriched in molecular chaperone genes, such as *HSP42*, *SSA4*, *HSP78*, *HSP82*, *HSP104*, and *SSE2*. Although a specific role for these chaperones in mating is not known, it is possible that elevated levels would interfere with mating. Assessing chaperone regulation during mating has been complicated by the difficulty of ensuring constant stress over the course of the experiment. Indeed, different studies have yielded conflicting data (39, 46). However, the Kar4 dependence of their regulation suggests that repression is real.

Cluster 6 is comprised of genes showing similar levels of expression during vegetative growth and after pheromone treatment in wild type but which showed significant induction by pheromone in the *kar4Δ* strain. Cluster 6 was enriched in genes known to play a role in the mating response, encoding proteins involved in pheromone synthesis (*AXL1*, *MFα1*, and *MFα2*), pheromone receptors (*STE3*), agglutination (*SAG1* and *MUC1*), and signaling (*FUS3*). Most of these genes show relatively little change during an extended time course of pheromone response (39). Because most of these genes act in the earliest stages of the mating response, delayed Kar4-dependent negative regulation may allow for a pulse of expression preparatory to mating.

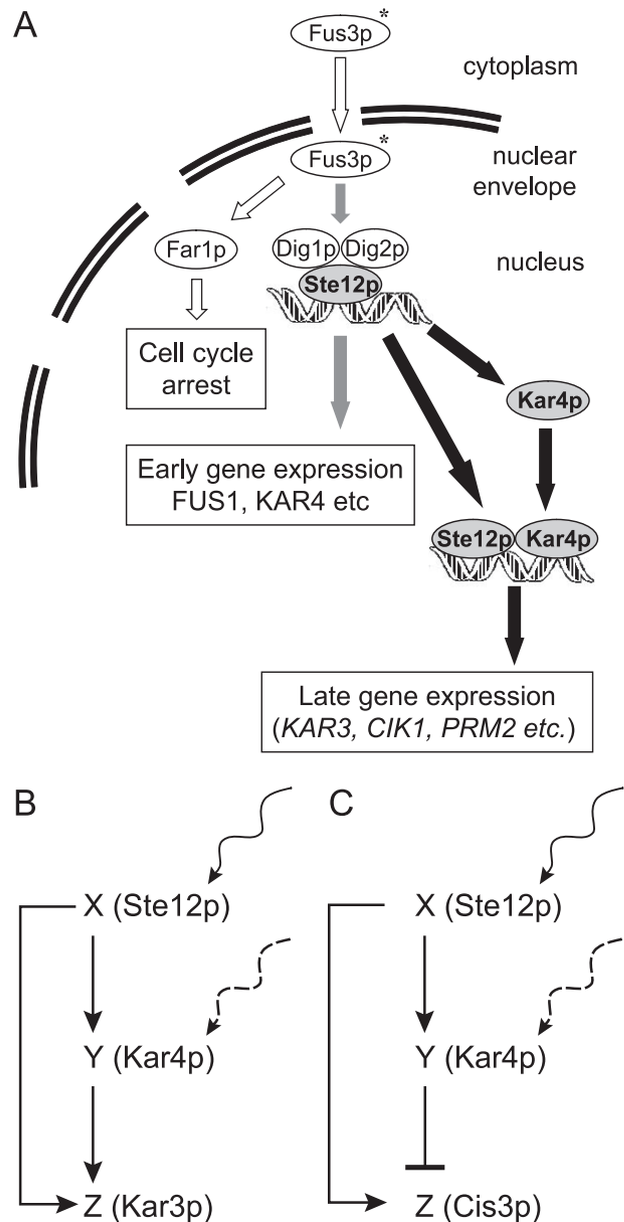


FIG. 8. (A) Proposed model of Kar4 action in the pheromone response. During the pheromone response, cytoplasmic Fus3 (Fus3-C) enters the nucleus (Fus3-N), where it phosphorylates Far1 and Dig1 and Dig2, leading to cell cycle arrest, and the activation of Ste12, which in turn induces the early pheromone response genes. Ste12 also induces the expression of the pheromone-specific short form of Kar4, which begins to accumulate in the cell. After sufficient Kar4 has accumulated, it facilitates Ste12 binding to the promoter regions of genes involved in functions required late in the pheromone response. (B) Representation of the feed-forward loop. The diagram was modified from that of Mangan and Alon (29). Coherent type I feed forward loop (AND gate logic) in which transcription factor X is represented by Ste12 and transcription factor Y is represented by Kar4 and the target gene, Z, is represented by *KAR3*. The inducer of X and Y is pheromone, although other inducers of Kar4 are possible. This type of feed-forward loop is suggested to act as a persistence detector where only persistent stimulus of both X and Y will lead to activation of Z (29). (C) Incoherent type I feed-forward loop (AND gate logic) in which X is represented by Ste12, Y is represented by Kar4, and Z is represented by *CIS3*. This type of feed-forward loop is characterized by the combined induction of X and Y by a stimulus (pheromone) in order to cause Z to be repressed by Y.

Genes in cluster 5 were also constitutively expressed in the wild-type strain but, in contrast to cluster 6, were downregulated by pheromone in the *kar4* Δ mutant. One hypothesis to explain cluster 5 is that it is comprised of genes whose expression is cell cycle regulated and not normally expressed during the G_1 phase. However, expression of these genes may be required during mating and therefore induced by pheromone. Consistent with this, cluster 5 is enriched in genes that are involved in RNA processing and ribosome biogenesis, and may facilitate high levels of protein synthesis during mating. Several genes within this cluster (*RPB11*, *MAK16*, *TRM1*, *SRP40*, *NSR1*, *RRP8*, and *HCA4*) have been shown to be expressed in pheromone-arrested cells, repressed as the cells reenter the cell cycle at G_1 , and then induced later in the cell cycle (46).

Cluster 7 was comprised of genes that are expressed constitutively in wild-type cells but showed strongly elevated levels of expression during mitotic growth in the *kar4* Δ mutant. The behavior of these genes suggests that Kar4 may negatively regulate the expression of some genes during vegetative growth. Because their levels of gene expression are not elevated in the *kar4* Δ mutant after pheromone treatment, these genes must not be expressed at G_1 but would be expressed at other times in the cell cycle. Consistent with this hypothesis, cluster 7 is enriched in genes involved in cell cycle regulation (*CDC5* and *CDC6*), as well as in microtubule and spindle pole body function (*SPC42*, *VIK1*, and *SPC24*).

Temporal regulation of Kar4-dependent pheromone induced genes. The Kar4-dependent pheromone-induced genes in cluster 3 were enriched in genes induced later in the pheromone response. Because Kar4 is itself induced by pheromone, it was proposed that Kar4 may function as a temporal regulator of a subset of mating-specific proteins (23).

In support of this hypothesis, premature expression of Kar4 significantly accelerated the expression of *KAR3* and *PRM2*, causing both genes to become half-maximally induced within 5 min after pheromone addition. The accelerated kinetics match those observed for the bulk of the genes whose expression is independent of Kar4 (Fig. 6). In wild-type cells, *KAR3* expression was induced more quickly than *PRM2*, indicating that the various Kar4-dependent promoters may respond to different levels of Kar4.

These results suggest that, with respect to pheromone-dependent transcriptional induction, Kar4 acts like a type I coherent feed-forward loop (Fig. 8B) (29, 30). Briefly, a feed-forward loop is composed of a transcription factor X (e.g., Ste12) that regulates a second transcription factor Y (e.g., Kar4), such that both regulate downstream genes, Z (e.g., *KAR3* and other cluster 2 genes induced later in the pheromone response). In the AND-gate version of the type I coherent feed-forward loop, the requirement for the second transcription factor reaching a concentration threshold results in a sigmoid-shaped delayed response. This model predicts that constitutive expression of activating levels of Kar4 would transform the system into one exhibiting a simple dependence on the activating stimulus, erasing both the delay and the sigmoid behavior, as we observed (Fig. 7C).

With respect to genes showing Kar4-dependent repression on pheromone (clusters 4 and 6), a second class of a feed-forward loop may be more relevant. In the type I incoherent feed-forward loop with AND-gate logic (Fig. 8C), a transcrip-

tion factor X (i.e., Ste12) activates both gene Z (e.g., *Cis3*) and a second transcription factor Y (i.e., Kar4), which in turn represses gene Z. *CIS3* expression is repressed during pheromone response in both a Ste12 and Kar4-dependent manner (our data and reference 39), although the dependence on Ste12 may simply reflect the requirement for Kar4 expression. One characteristic of incoherent type I feed-forward loops is that they show a modest pulse of Z expression upon stimulation of X, since it takes time for Y to accumulate. Cluster 6 included several genes that act early in mating and so might be expected to be expressed at a high level early, but not later, in the mating response.

Kar4 regulatory motif. Previous experiments demonstrated the Kar4-dependent transcriptional regulation of *KAR3* and *CIK1* but did not establish that Kar4 binds to DNA. Given the sequence homology between Kar4 and mRNA methyltransferases, it was critical to reexamine whether Kar4 interacts with DNA. In this work, we showed that Kar4 is associated with the chromatin upstream of *KAR3* and that purified Kar4 binds to DNA regulatory sequences upstream of *KAR3* in vitro. Furthermore, Kar4-dependent regulation was observed using a heterologous reporter construct (*CYC1-lacZ*) containing the upstream regulatory elements but no mRNA sequences from *KAR3* (23; the present study). Thus, Kar4-dependent regulation during mating most likely acts via regulatory DNA sequences rather than by elements present within the mRNA.

To characterize the Kar4 regulatory sequence, we used AlignAce (18) to identify DNA motifs enriched upstream of the genes showing strong Kar4-dependent pheromone induction. This analysis identified a motif, C/TAANNCAAANN CN GNYT, contained within the minimal region of *KAR3*, which was necessary for pheromone induction. Interestingly, this motif was not enriched in the other clusters showing Kar4-dependent regulation. Kar4 may either (i) bind to more than one motif, (ii) bind via interactions with a second DNA-binding protein, or (iii) act indirectly through the induction of other regulatory proteins. Motif analysis yielded a large set of DNA motifs enriched in the upstream regulatory regions of the other clusters, but it remains to be determined whether any are required for Kar4-dependent regulation.

In sum, we have shown that Kar4 plays a more extensive and complex role in yeast mating than previously thought. Interestingly, the human pathogen *Candida albicans* contains a homolog of Kar4 that is transcriptionally regulated both by mating pheromone and by whether the cell is in the alternate "opaque" phase of growth (24). Similar to our observations with budding yeast, several opaque-phase genes, including gene associated with skin colonization are downregulated during the pheromone response. These results suggest that in *Candida*, as in *Saccharomyces*, Kar4 may help provide part of the complex pattern of gene regulation during different cell developmental phases of growth.

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REFERENCES

1. Ausubel, F. M. 1987. Current protocols in molecular biology. Greene Publishing Associates, Media, PA.

2. Baur, M., R. K. Esch, and B. Errede. 1997. Cooperative binding interactions required for function of the Ty1 sterile responsive element. *Mol. Cell. Biol.* **17**:4330–4337.
3. Beh, C. T., V. Brizzio, and M. D. Rose. 1997. *KAR5* encodes a novel pheromone-inducible protein required for homotypic nuclear fusion. *J. Cell Biol.* **139**:1063–1076.
4. Bohm, S., D. Frishman, and H. W. Mewes. 1997. Variations of the C2H2 zinc finger motif in the yeast genome and classification of yeast zinc finger proteins. *Nucleic Acids Res.* **25**:2464–2469.
5. Bujnicki, J. M., M. Feder, M. Radlinska, and R. M. Blumenthal. 2002. Structure prediction and phylogenetic analysis of a functionally diverse family of proteins homologous to the MT-A70 subunit of the human mRNA: m⁶A methyltransferase. *J. Mol. Evol.* **55**:431–444.
6. Burke, D., D. Dawson, T. Stearns, et al. 2000. *Methods in yeast genetics*. Cold Spring Harbor Laboratory Press, Plainview, NY.
7. Clancy, M. J., M. E. Shambaugh, C. S. Timpte, and J. A. Bokar. 2002. Induction of sporulation in *Saccharomyces cerevisiae* leads to the formation of N⁶-methyladenosine in mRNA: a potential mechanism for the activity of the IME4 gene. *Nucleic Acids Res.* **30**:4509–4518.
8. Desjarlais, J. R., and J. M. Berg. 1992. Toward rules relating zinc finger protein sequences and DNA binding site preferences. *Proc. Natl. Acad. Sci. USA* **89**:7345–7349.
9. Dohlman, H. G., and J. W. Thorner. 2001. Regulation of G protein-initiated signal transduction in yeast: paradigms and principles. *Annu. Rev. Biochem.* **70**:703–754.
10. Dolan, J. W., C. Kirkman, and S. Fields. 1989. The yeast STE12 protein binds to the DNA sequence mediating pheromone induction. *Proc. Natl. Acad. Sci. USA* **86**:5703–5707.
11. Elion, E. A., M. Qi, and W. Chen. 2005. Signal transduction: signaling specificity in yeast. *Science* **307**:687–688.
12. Errede, B., and G. Ammerer. 1989. STE12, a protein involved in cell-type-specific transcription and signal transduction in yeast, is part of protein-DNA complexes. *Genes Dev.* **3**:1349–1361.
13. Fitch, P. G., A. E. Gammie, D. J. Lee, V. B. de Candal, and M. D. Rose. 2004. Lrg1p Is a Rho1 GTPase-activating protein required for efficient cell fusion in yeast. *Genetics* **168**:733–746.
14. Gammie, A. E., B. G. Stewart, C. F. Scott, and M. D. Rose. 1999. The two forms of karyogamy transcription factor Kar4p are regulated by differential initiation of transcription, translation, and protein turnover. *Mol. Cell. Biol.* **19**:817–825.
15. Guarente, L., and M. Ptashne. 1981. Fusion of *Escherichia coli lacZ* to the cytochrome c gene of *Saccharomyces cerevisiae*. *Proc. Natl. Acad. Sci. USA* **78**:2199–2203.
16. Hagen, D. C., G. McCaffrey, and G. F. Sprague, Jr. 1991. Pheromone response elements are necessary and sufficient for basal and pheromone-induced transcription of the *FUS1* gene of *Saccharomyces cerevisiae*. *Mol. Cell. Biol.* **11**:2952–2961.
17. Heiman, M. G., and P. Walter. 2000. Prm1p, a pheromone-regulated multispansing membrane protein, facilitates plasma membrane fusion during yeast mating. *J. Cell Biol.* **151**:719–730.
18. Hughes, J. D., P. W. Estep, S. Tavazoie, and G. M. Church. 2000. Computational identification of cis-regulatory elements associated with groups of functionally related genes in *Saccharomyces cerevisiae*. *J. Mol. Biol.* **296**:1205–1214.
19. James, P., J. Halladay, and E. A. Craig. 1996. Genomic libraries and a host strain designed for highly efficient two-hybrid selection in yeast. *Genetics* **144**:1425–1436.
20. Kim, T. S., H. Y. Kim, J. H. Yoon, and H. S. Kang. 2004. Recruitment of the Swi/Snf complex by Ste12-Tec1 promotes Flo8-Mss11-mediated activation of *STB1* expression. *Mol. Cell. Biol.* **24**:9542–9556.
21. Kurdستاني, S. K., D. Robyr, S. Tavazoie, and M. Grunstein. 2002. Genome-wide binding map of the histone deacetylase Rpd3 in yeast. *Nat. Genet.* **31**:248–254.
22. Kurihara, L. J., C. T. Beh, M. Latterich, R. Schekman, and M. D. Rose. 1994. Nuclear congression and membrane fusion: two distinct events in the yeast karyogamy pathway. *J. Cell Biol.* **126**:911–923.
23. Kurihara, L. J., B. G. Stewart, A. E. Gammie, and M. D. Rose. 1996. Kar4p, a karyogamy-specific component of the yeast pheromone response pathway. *Mol. Cell. Biol.* **16**:3990–4002.
24. Lockhart, S. R., R. Zhao, K. J. Daniels, and D. R. Soll. 2003. Alpha-pheromone-induced “shmooing” and gene regulation require white-opaque switching during *Candida albicans* mating. *Eukaryot. Cell* **2**:847–855.
25. Mackey, A. T., L. R. Sproul, C. A. Sontag, L. L. Satterwhite, J. J. Correia, and S. P. Gilbert. 2004. Mechanistic analysis of the *Saccharomyces cerevisiae* kinesin Kar3. *J. Biol. Chem.* **279**:51354–51361.
26. Maddox, P., E. Chin, A. Mallavarapu, E. Yeh, E. D. Salmon, and K. Bloom. 1999. Microtubule dynamics from mating through the first zygotic division in the budding yeast *Saccharomyces cerevisiae*. *J. Cell Biol.* **144**:977–987.
27. Madhani, H. D., and G. R. Fink. 1997. Combinatorial control required for the specificity of yeast MAPK signaling. *Science* **275**:1314–1317.
28. Madhani, H. D., T. Galitski, E. S. Lander, and G. R. Fink. 1999. Effectors of a developmental mitogen-activated protein kinase cascade revealed by expression signatures of signaling mutants. *Proc. Natl. Acad. Sci. USA* **96**:12530–12535.
29. Mangan, S., and U. Alon. 2003. Structure and function of the feed-forward loop network motif. *Proc. Natl. Acad. Sci. USA* **100**:11980–11985.
30. Mangan, S., A. Zaslaver, and U. Alon. 2003. The coherent feed-forward loop serves as a sign-sensitive delay element in transcription networks. *J. Mol. Biol.* **334**:197–204.
31. Marsh, L., and M. D. Rose. 1997. The pathway of cell and nuclear fusion during mating in *Saccharomyces cerevisiae*, p. 827–888. In J. Pringle, J. R. Broach, and E. W. Jones (ed.), *The molecular and cellular biology of the yeast Saccharomyces cerevisiae*, vol. 3. Cell cycle and cell biology. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY.
32. McCaffrey, G., F. J. Clay, K. Kelsay, and G. F. Sprague, Jr. 1987. Identification and regulation of a gene required for cell fusion during mating of the yeast *Saccharomyces cerevisiae*. *Mol. Cell. Biol.* **7**:2680–2690.
33. Meluh, P. B., and M. D. Rose. 1990. *KAR3*, a kinesin-related gene required for yeast nuclear fusion. *Cell* **60**:1029–1041.
34. Miller, R. K., and M. D. Rose. 1998. Kar9p is a novel cortical protein required for cytoplasmic microtubule orientation in yeast. *J. Cell Biol.* **140**:377–390.
35. Oehlen, L. J., J. D. McKinney, and F. R. Cross. 1996. Ste12 and Mcm1 regulate cell cycle-dependent transcription of *FAR1*. *Mol. Cell. Biol.* **16**:2830–2837.
36. Ohashi, A., J. Gibson, I. Gregor, and G. Schatz. 1982. Import of proteins into mitochondria: the precursor of cytochrome c₁ is processed in two steps, one of them heme-dependent. *J. Biol. Chem.* **257**:13042–13047.
37. Oldenburg, K. R., K. T. Vo, S. Michaelis, and C. Paddon. 1997. Recombination-mediated PCR-directed plasmid construction in vivo in yeast. *Nucleic Acids Res.* **25**:451–452.
38. Page, B. D., L. L. Satterwhite, M. D. Rose, and M. Snyder. 1994. Localization of the Kar3 kinesin heavy chain-related protein requires the Cik1 interacting protein. *J. Cell Biol.* **124**:507–519.
39. Roberts, C. J., B. Nelson, M. J. Marton, R. Stoughton, M. R. Meyer, H. A. Bennett, Y. D. He, H. Dai, W. L. Walker, T. R. Hughes, M. Tyers, C. Boone, and S. H. Friend. 2000. Signaling and circuitry of multiple MAPK pathways revealed by a matrix of global gene expression profiles. *Science* **287**:873–880.
40. Robinson, M. D., J. Griggall, N. Mohammad, and T. R. Hughes. 2002. FunSpec: a web-based cluster interpreter for yeast. *BMC Bioinformatics* **3**:35–39.
41. Rose, M. D. 1996. Nuclear fusion in the yeast *Saccharomyces cerevisiae*. *Annu. Rev. Cell Dev. Biol.* **12**:663–695.
42. Rose, M. D., F. M. Winston, P. Hieter, et al. 1990. *Methods in yeast genetics: a laboratory course manual*. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY.
43. Sambrook, J., E. F. Fritsch, and T. Maniatis. 1989. *Molecular cloning: a laboratory manual*, 2nd ed. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY.
44. Schmitt, M. E., T. A. Brown, and B. L. Trumppower. 1990. A rapid and simple method for preparation of RNA from *Saccharomyces cerevisiae*. *Nucleic Acids Res.* **18**:3091–3092.
45. Sikorski, R. S., and P. Hieter. 1989. A system of shuttle vectors and yeast host strains designed for efficient manipulation of DNA in *Saccharomyces cerevisiae*. *Genetics* **122**:19–27.
46. Spellman, P. T., G. Sherlock, M. Q. Zhang, V. R. Iyer, K. Anders, M. B. Eisen, P. O. Brown, D. Botstein, and B. Futcher. 1998. Comprehensive identification of cell cycle-regulated genes of the yeast *Saccharomyces cerevisiae* by microarray hybridization. *Mol. Biol. Cell* **9**:3273–3297.
47. Sprague, G. F., and J. W. Thorner. 1992. Pheromone response and signal transduction during the mating process of *Saccharomyces cerevisiae*, p. 657–744. In J. R. Broach, J. Pringle, and E. W. Jones (ed.), *The molecular and cellular biology of the yeast Saccharomyces cerevisiae*, vol. 2. Gene expression. Cold Spring Harbor Laboratory Press, Cold Spring Harbor, NY.
48. Tavazoie, S., J. D. Hughes, M. J. Campbell, R. J. Cho, and G. M. Church. 1999. Systematic determination of genetic network architecture. *Nat. Genet.* **22**:281–285.
49. Winzler, E. A., D. D. Shoemaker, A. Astromoff, H. Liang, K. Anderson, B. Andre, R. Bangham, R. Benito, J. D. Boeke, H. Bussey, A. M. Chu, C. Connelly, K. Davis, F. Dietrich, S. W. Dow, M. El Bakkoury, F. Foury, S. H. Friend, E. Gentalen, G. Gaever, J. H. Hegemann, T. Jones, M. Laub, H. Liao, N. Liebunghuth, D. J. Lockhart, A. Lucau-Danila, M. Lussier, L. M'Rabet, P. Menard, M. Mittmann, C. Pai, C. Rebischung, J. L. Revuelta, N. Riles, C. J. Roberts, P. Ross-MacDonald, B. Scherens, M. Snyder, S. Sookhai-Mahadeo, R. K. Storms, S. Veronneau, M. Voet, G. Volckaert, T. R. Ward, R. Wysocki, G. S. Yen, K. Yu, K. Zimmermann, P. Philippsen, M. Johnston, and R. W. Davis. 1999. Functional characterization of the *Saccharomyces cerevisiae* genome by gene deletion and parallel analysis. *Science* **285**:901–906.
50. Zeitlinger, J., I. Simon, C. T. Harbison, N. M. Hannett, T. L. Volkert, G. R. Fink, and R. A. Young. 2003. Program-specific distribution of a transcription factor dependent on partner transcription factor and MAPK signaling. *Cell* **113**:395–404.